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14. ABSTRACT <p>This report results from a contract tasking Fraunhofer Institut für Chemische Technologie as follows: Lattice defects in energetic materials will be measured using X-ray diffraction. The method shall be tested for its ability to characterize lattice defects in RDX and HMX, where dislocations gliding and deformation twinning are believed to dominate mechanical behavior. X-ray diffraction patterns will be evaluated relating to phase, residual strain, crystallite size and micro strain. The occurring phases and the lattice parameters of the samples will be determined by Rietveld-method using literature data. Micro strain and crystallite size will be determined with the Williamson Hall method by plotting reciprocal peak widths versus reciprocal lattice distances. New routines for size/strain evaluation, implemented in Rietveld programs will be tested and compared with the Williamson Hall method.</p>					
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Characterization of the Microstructure of fine Energetic Materials

Final Report

by
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Summary

Mechanical sensitivity of energetic materials as explosives and propellants has been discussed as a function of particle size and lattice defects originated during crystallization and particle processing. As a conclusive correlation hitherto is difficult to achieve an approach is made to investigate lattice defects using X-ray diffraction.

SEM and X-ray diffraction measurements were carried out with fine RDX and HMX. These materials have the interesting feature that different defect mechanisms are supposed to dominate the mechanical behavior. The X-ray diffraction patterns were evaluated by Rietveld- and peak profile analysis. The X-ray systems has been calibrated using the profile standard SRM 660a from the National Institute for Standards and Technology (NIST).

A new preparation technique was tested in order to reduce broadening due to the low absorption, particularly in energetic materials like nitramines.

The investigations reveal interesting details about lattice defects in energetic materials. Characteristic peak broadening was found for the nitramines RDX and HMX. The results agree with the models described in literature, pointing to a higher micro strain in HMX. The two-dimensional interfaces of twins in HMX seem to create higher micro strain then the one dimensional dislocations in RDX. Besides twin interfaces may divide crystallites into smaller coherently diffracting domains. Moreover an unisotropy peak broadening was found for RDX, which were discussed in terms of strain fields created by unisotropic dislocation.

List of keywords

Energetic materials, lattice defects, micro strain, X-ray diffraction

Introduction

Mechanical sensitivity is an important issue for energetic materials as explosives and propellants. It has been discussed as a function of particle size and lattice defects originated during crystallization and particle processing. However, a conclusive correlation was hitherto difficult to achieve, because the integral measurement of defects is difficult. An approach is made to quantify and characterize lattice defects in energetic materials by means of X-ray diffraction.

State of the art

X-ray diffraction

The X-ray diffraction allows a characterization of energetic crystals and particles obtained not only from the surface and with point measurements but from a representative volume of a substance. The method is based on the evaluation of diffraction line profiles; their positions, intensities, widths and shapes. The line intensities and positions are generally used to identify and quantify crystalline ingredients or phases, to determine or refine crystal structures or investigate residual strain.

The line widths have many physical sources. The measuring system reveals a geometrical profile representing optical devices and X-ray tube. This profile is broadened by sample features as size, strain, diffuse scattering, misfits between phases, etc. The effects superimpose and often are not easy to separate. Crystal size and lattice distortion broadening are usual present concomitantly, and strain broadening can be caused by many different crystal defects as dislocations, stacking faults, point defects, misfits, distorted grain boundaries, concentration gradients, etc. The Warren-Averbach-method solves the problem of the separation of size and strain broadening revealing the root mean-squared strain ($\langle \epsilon^2 \rangle^{1/2}$). (Ungar et al. 2000) Williamson and Hall (1953) suggested that the full width at half-maximum (FWHM) of diffraction profiles can be written as the sum of strain and size broadening, which means in terms of reciprocal units: $\beta^* = \Delta K = 0.9/D + \Delta K^D$ where ΔK^D is the strain contribution to peak broadening and D is the average grain size. ($d^* = K = 2 \lambda \sin \theta$, $\Delta K = 2 \Delta \theta / \lambda \cos \theta$ with the peak width β , the lattice spacing d , the diffraction angle θ , the half of FWHM $\Delta \theta$ and the wave length λ)

A crucial point in this method is the physical model placed behind the root-mean-squared strains. If the displacement of atoms or molecules relative to their ideal position is random the root-mean-squared strain is a constant. In most cases, however, strain is caused by dislocations, and in this case the displacements have a well defined spatial distribution and the root-mean-squared strain is far from being constant. Recent results obtained by analyzing strain anisotropy enabled a microscopic model for the root-mean-squared strains based on dislocation theory, called *modified* Warren-Averbach analysis and *modified* Williamson-Hall plot. (Ungar et al. 2000)

Beside the Williamson-Hall method, where strain and size broadening are separated on the basis that size or strain is diffraction order independent or dependent, other methods bases on a profile analysis of single peaks, as the shapes of size and strain components are different (e.g. Manual of TOPAS from Bruker AXS).

For calibration and determination of geometrical profiles powder diffraction standard reference materials are available from the National Institute of Standards and Technology NIST. The line position and line shape standard SRM 660a is especially recommended for size-strain investigations as it is more or less strain-free and yields no size broadening. It consists of a fine Lanthanum Hexaboride powder with aggregates of crystallizes in the 2 to 5 μm range. (NIST 2000).

Energetics - actual research and structures

According to the discussions mentioned in the introduction concepts were developed to reduce defect concentrations of energetic materials applying careful crystallization and processing methods. Particularly, energetic nitramines as HMX and RDX are in the scope of interest, and for these materials it was shown that an improved internal crystal quality decreases the mechanical sensitivity.

In case of RDX an insensitive industrial product "I-RDX" is already available from SNPE, France. (Bouma et al. 2001, Lécume et al. 2001, Kröber et al. 1998, Van der Heijden et al. 1998).

In order to explain these effects special lattice imperfections are discussed to create hot spots under mechanical stress (Armstrong et al. 1993), and detailed models of embedded defects as dislocations and twins were developed.

Crystal and real structure of RDX ($C_3H_6N_6O_6$)

The crystal structure of RDX has been refined from single-crystal neutron-diffraction data by Choi and Prince 1972. The compound crystallizes in the orthorhombic space group $Pbca$ with the lattice parameters $a=13.182$, $b=11.574$ and $c=10.709$ Å. The elementary cell contains eight RDX molecules, measured and calculated densities amount to 1,816 and 1,806, respectively.

Armstrong and Elban described that the mechanical properties of RDX are explained by the difficulty of moving individual dislocations. Fig. 1 shows the $[-100](040)$ edge dislocation of the RDX lattice, schematically (Armstrong et al.). The dislocation was constructed to explain the (010) slip reported by Cinnick and May in 1969. Later on Elban, Armstrong and colleagues reported the occurrence of slip on (021) and (02-1) systems in RDX.

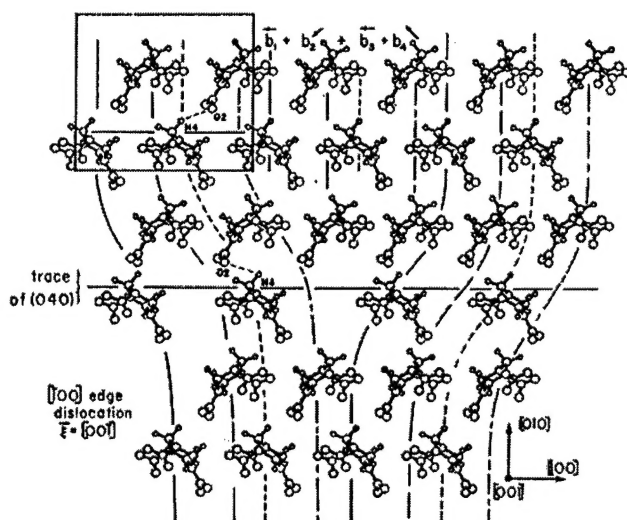


Fig. 1: $[-100](040)$ edge dislocation in RDX, schematically

Crystal and real structure of HMX ($C_4H_8N_8O_8$)

HMX crystallizes in four phases; α -, β -, γ - and δ -HMX. β -HMX is the most interesting phase as it is the room temperature stable phase with the highest density. Eiland and Pepinsky found in 1955 space group $P21/n$ with lattice parameters $a=6.54$, $b=11.05$, $c=7.37$ Å and $\beta=102.8^\circ$. Reinvestigations by Choi and Boutin in 1970 revealed space group $P21/c$ with the lattice parameters $a=6.54$, $b=11.05$, $c=8.70$ Å and $\beta=124.3^\circ$. The elementary cell contains two molecules, measured and calculated densities amount to 1.894 and 1.90 g/cm³, respectively.

The internal properties of β -HMX seem to be dominated by crystal twinning. Palmer et al. 1982, Cady 1993 and Armstrong et al. 1993 observed (101)-twins in HMX. The twins was found in crystals grown from solutions and when crystals are exhibited to mechanical stress (deformation twinning), and the deformation twinning was found to be reversible at low stress. The twin system is characterized by the specifications $K_1, \eta_1, K_2, \eta_2 = (101), [10-1], (10-1), [101]$. It has been reasoned to be of type II in which the twinning orientation relationship may be visualized to occur by a rotation of 180° about the twinning η_1 direction, followed by a parallel translation. (Armstrong et al. 1993)

Objectives and approach

A first approach was made by Engel and Herrmann (2001) with the seed project "Lattice Imperfections of Energetic Materials Measured with X-ray Diffraction" supported by the U.S. Government through its European Research Office of the U.S. Army. Measurements were carried out with relatively large HMX particles crystallized from solutions. The investigations revealed that the method is capable for detecting qualities of coarse crystals, when suited measuring systems are applied to reduce effects of uneven sample surfaces and orientation, and after all show a correlation of micro strain with sensitivity against friction.

With the new project the quantification of micro strain using X-ray diffraction shall be verified especially for fine materials, where effects of uneven surfaces and badly distributed orientations do not appear. Since diffraction peak broadening occurs with small crystallites below a few μm , superimposing with broadening by micro strain, samples with a particle or crystallite size between 20 and one μm shall be used. The characterization of the internal quality shall include the occurring phases, crystallite and particle size, lattice parameters and mean squared micro strain. Particular interest is drawn to the characterization of defect structures. Therefore the unisotropy of micro strain and particle size in RDX and HMX shall be determined and compared with the lattice models described in literature and their orientation.

Experimental

For the investigations comminuted powders of HMX and RDX with particle sizes of a few microns were provided by the particle processing group of ICT, and samples of the fine powders were investigated under the scanning electron microscopy (SEM) for a first characterization of the crystal quality and morphology.

Diffraction patterns were measured with a diffractometer D5000 from Bruker AXS, equipped with copper tube, vertical soller slits, Ni- $k\beta$ -filter, divergence and detector slits of $0,3^\circ$ and $0,1\text{ mm}$, and rotating sample holder. Tab. 1 summarizes the instrumental parameters and Tab. 7 (Appendix) the diffraction measurements and their conditions. The measurements are designated by X followed by a number, e.g. X3 is the third diffraction measurement with file name mh091203. The measurements will be referred by these numbers in the following chapters. For determining the geometrical profiles of the systems the standard reference material SRM 660a (NIST 2000) was measured under the same conditions as the energetic materials.

A preliminary series was measured with common sample preparation, where the powders were filled up to the edge of a standard sample holder and the surfaces were flattened with a glass plate (X1-X4). Patterns were measured between 5 and $90^\circ 2\theta$ with a step width of $0.01^\circ 2\theta$ and a measuring time of 8 s/step ($\sim 19\text{ h/pattern}$).

Tab. 1: Instrumental parameters

Goniometer Radii	Value	Full Axial Model	Value
Primary Radius (mm)	217.5	Source Length (mm)	12
Secondary Radius (mm)	217.5	...Source Width (mm)	0.04
Equatorial Convolutions		Sample Length (mm)	14
RS Width (mm)	0.1	RS Length (mm)	12
FDS Angle ($^\circ$)	$0.3 / 0.5$	Prim. Soller ($^\circ$)	2.3
		Sec. Soller ($^\circ$)	2.3

As strong absorption broadening was found with these measurements, particularly with the energetic samples, a second series was started, preparing samples in capillaries on a Debye-Scherrer device (X5-X8), and as thin powder layers prepared on a wafer or an aluminum plate (X9-X16), in order to reduce absorption broadening by reducing the diffracting volume in the sample. The patterns were measured between 10 and 40 °2Theta.

Refined patterns of the powder layers were measured in the reduced 2Theta ranges between 12.5 and 26 °2Theta for RDX, and 14 and 33 °2Theta for HMX and LaB6. Each sample was measured twice, freshly prepared, with 0.005 2Theta step width and 10 s measuring time per step.

Evaluation

The diffraction patterns measured with the common sample preparation were evaluated using the full pattern fitting tools of the program TOPAS from Bruker/AXS, refining peak positions of so-called hkl-phases calculated from crystal data reported in literature. Peaks were fitted using the ray tracing fundamental parameter approach (FPA), deriving peak profiles from the optical arrangement of the measuring system. Thus phases were identified, peaks were indexed, lattice parameters were refined, and peaks with minimal or without overlap were identified as »single peaks«.

The single peaks were fitted using the split-Pearson VII (SPVII) analytical function and the Fundamental Parameter Approach FPA of program TOPAS.

The evaluation with SPVII yields Full Widths at Half Maximum (FWHM) of the peaks, which were reduced by the geometric peak widths obtained from the standard measurements, in order to obtain pure sample broadening. The data were evaluated in terms of reciprocal units as described by Williamson and Hall. Therefore reciprocal peak widths β^* were plotted versus reciprocal lattice distances d^* and linear regression lines were fitted in the low d^* -range. The slopes and intercepts of linear regression lines with the y-axis are discussed in terms of micro strain and inverse particle size. Additionally, deviations of the reciprocal peak widths from regression lines were determined and related to their peak indices, and systematic deviations were compared with unisotropic defect models reported in literature.

For the evaluation with the FPA geometrical profiles were parameterized on the basis of the SRM 660a pattern, and the three strongest peaks of each energetic materials pattern were evaluated, revealing a main micro strain and particle size.

The linear absorption coefficients were calculated as described by Klug and Alexander 1974 for an estimation of peak broadening due to low absorption.

Results

Particle size and morphology (SEM)

Fig. 2 and Fig. 3 show selected pictures of RDX and HMX obtained with scanning electron microscopy. Both samples consist of particles in the μm -scale. Medium particle sizes may lie between 3 and 5 μm . Comparing the samples, the size distribution of the RDX-particles is narrower, and the surfaces of the RDX-particles are smoother and more rounded than those of the HMX-particles.

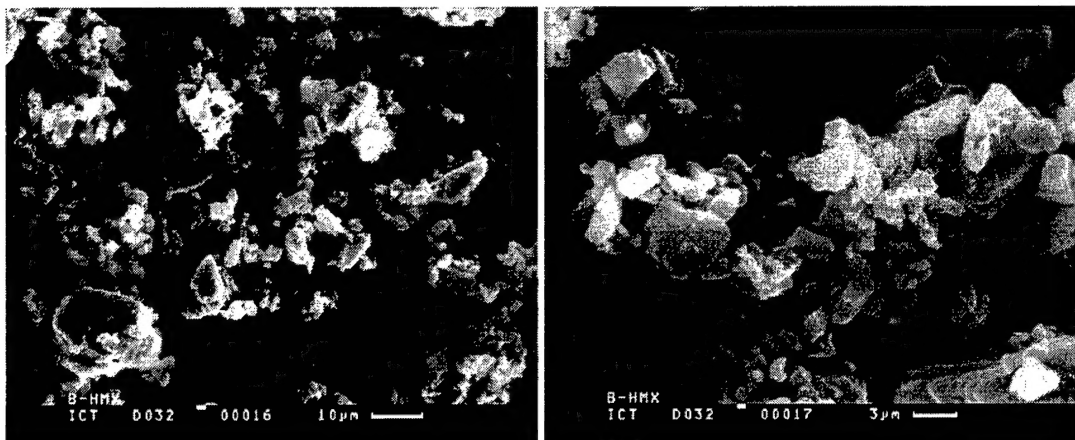


Fig. 2: Selected SEM-pictures of HMX-particles.

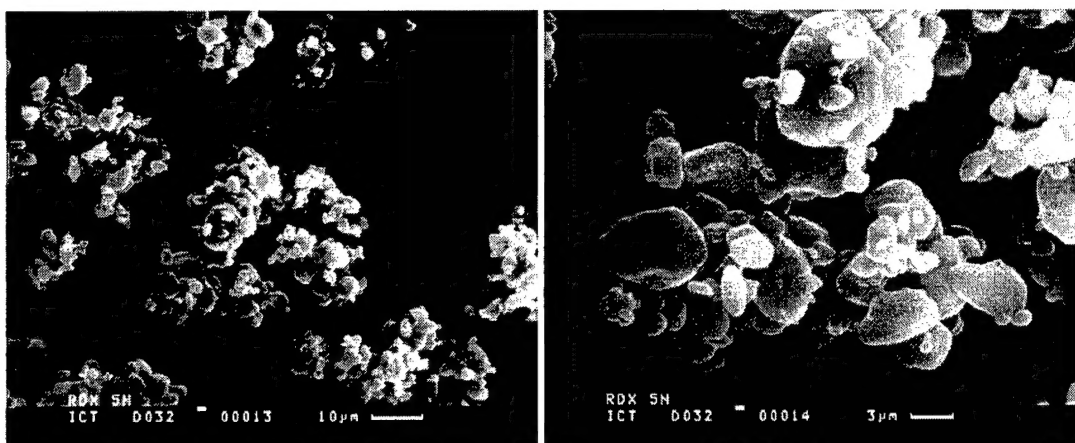


Fig. 3: Selected SEM-pictures of RDX-particles.

Linear absorption coefficients

The densities, linear absorption coefficients and penetration depths calculated according to Klug and Alexander 1974 are summarized in Tab. 2. The reference material has a much higher absorption coefficient than the energetic materials, resulting in penetration depth of 16 µm for LaB₆ and 642 and 614 µm for RDX and HMX, respectively.

Tab. 2: Absorption coefficients and penetration depths for X-ray radiation (CuKα)

	Density [g/cm ³]	Lin. absorption coeff. [1/cm]	Penetration depth [µm]
RDX	1.82	15.6	642
HMX	1.9	19.3	614
LaB ₆ (SRM 660a)	4.711 (ICDD 34-427)	1098.7	9.1

Comparison of measuring techniques

Fig. 4 shows patterns of RDX measured with the Debye-Scherrer device with rotating capillary (X5), and with Bragg-Brentano geometry with common sample preparation (X2) and with powder layers prepared on an aluminum plate (X14). Resulting peak widths for peaks (111) RDX at 13,1 2Theta and (100) LaB₆ at 21.35 °2Theta are summarized in Tab. 3.

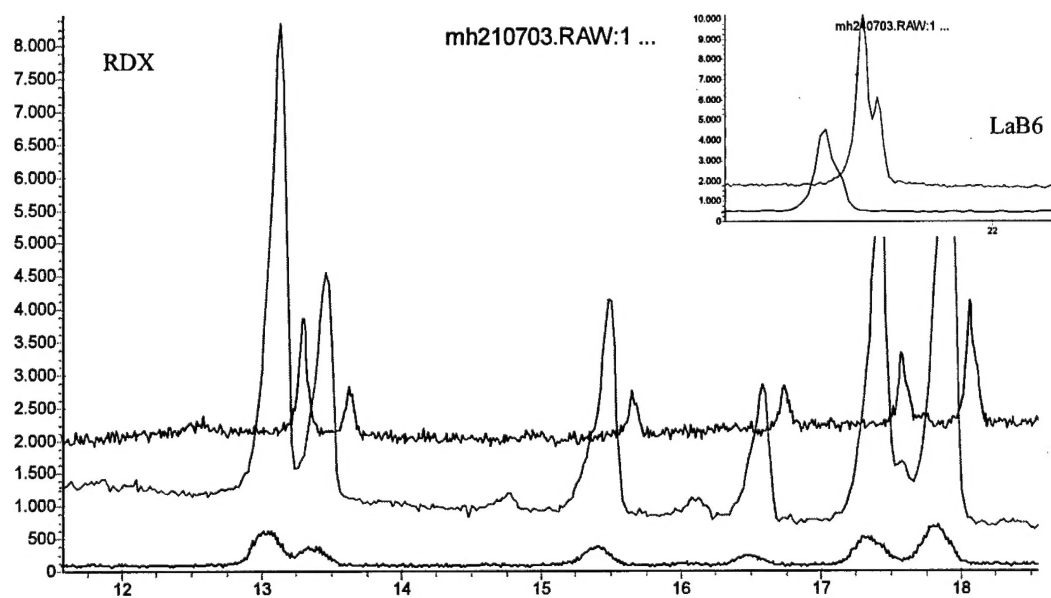


Fig. 4: Diffraction patterns of RDX measured with Debye-Scherrer device (black), Bragg-Brentano with common sample preparation (green) and powder layers on aluminum plate (blue), and of LaB₆ (small diagram) measured with common sample preparation (black) and powder layer (blue).

Tab. 3: Peak widths FWHM measured with different systems and preparation techniques

	LaB ₆	RDX
	(100) at 21.41 °2θ	(111) at 13.07 °2θ
Debye-Scherrer		0.198
Bragg-Brentano	0.054	0.123
Bragg-B., Layer	0.037	0.053

The Debye-Scherrer measurements yielded relatively broad peaks widths, e.g. 0.198 °2Theta for (111)RDX, with a low intensity. Narrower peaks were measured with the Bragg-Brentano geometry with 0.123 °2Theta of (111)RDX, and 0.054 °2Theta of (100)LaB₆. The narrowest peaks were obtained with the powder layers prepared on an aluminum plate with peak widths of 0.053 of (111)RDX, and even 0.037 °2Theta of (100)LaB₆. However, the patterns show also that for the high resolution obtained from powder layers poor peak intensities has to be accepted. Additionally, peak shifts indicate a poor adjustment of the sample position, with the already makeshift preparation technique.

The patterns with the highest intensities obtained with common preparation technique are used for the Rietveld-analysis. For the investigation of peak broadening the measurements with powder layers seem to provide the best basis, if the lack of intensity is compensated by a longer measuring time per step and/or a reduced step size. Besides, selected 2Theta ranges may be measured to limit the required

measuring time per pattern. Fig. 5 shows for instance a pattern X19 (blue) measured with a measuring time of 10 s/0.005 °2Theta. The intensity to underground ratio of this pattern is much better than with the measurement X14 (green) with the shorter measuring time and larger step width.

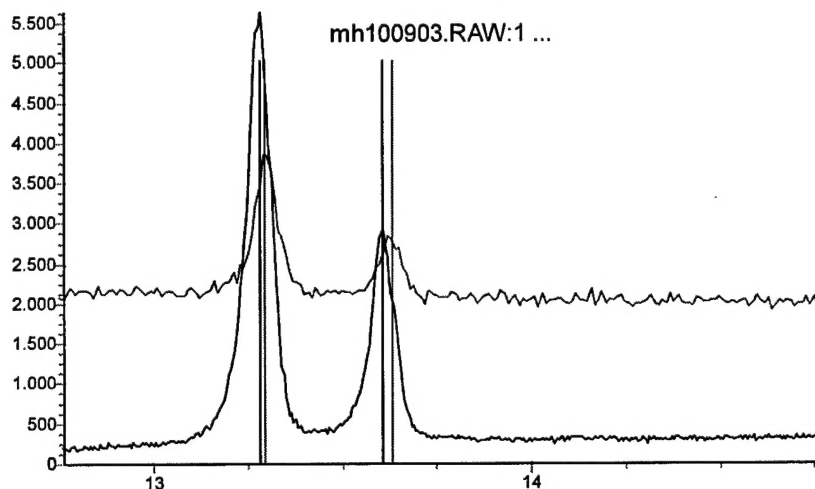


Fig. 5: Diffraction patterns of RDX measured with powder layers and different measuring times and step widths.

Full pattern fit (XRD-Bragg-Brentano/Rietveld)

Fig. 6 und Fig. 7 show the diffraction patterns X2 of RDX and X1 of HMX and their full pattern fits. All measured peaks belong to the structures of RDX and beta-HMX. However, the RDX-sample contains also HMX indicated by peaks at 14.7 and 16.2 °2Theta.

The refined lattice parameters are summarized in Tab. 4, peak positions and indices in Tab. 9 (Appendix). The Rietveld-analysis yielded good fits of RDX and HMX starting from the data reported by Choi et al. 1972 and Cady at al. 1963. Refined parameters are $a=13.205$, $b=11.616$ and $c=10.730$ Å and $\alpha=6.537$, $b=11.034$, $c=8.699$ Å and $\beta=124.44^\circ$ for RDX and HMX, respectively. The lattice parameters of RDX seem to be slightly strengthened compared to the literature values, whereas those of HMX agree well with the literature data.

Tab. 4: Lattice parameters obtained by Rietveld-refinement and from literature

Lattice parameters	RDX		HMX	
	Refinement	Choi et al. 1972	Refinement	Cady at al. 1963
a [Å]	13.205	13.182	6.537	6.54
b [Å]	11.616	11.574	11.034	11.05
c [Å]	10.730	10.709	8.699	8.70
β [°]			124.44	124.3

The peak positions and the indexing in Tab. 9 identify separated peaks of RDX at 13.07 (111), 13.40 (200), 16.51 (002), 17.35 (021), 17.83 (102), 20.35 (220), 21.99 (221), 25.35 (131), 30.42 (322), 31.14 (420), and of HMX at 14.70 (011), 18.28 (110), 20.53 (10-2), 22.06 (11-2), 23.02 (120), 27.19 (031), 29.64 (022), 31.90 (13-2), and 33.17 (200). Additionally the RDX-peak (210) at 14.42 °2Theta and the HMX-double peak (11-1)/(020) at 6.02 °2Theta may be used for the evaluation, taking into account

the overlaps. In case of the double peaks the resulting peak widths were reduced by the shift of $0.011^{\circ}2\theta$ between these peaks.

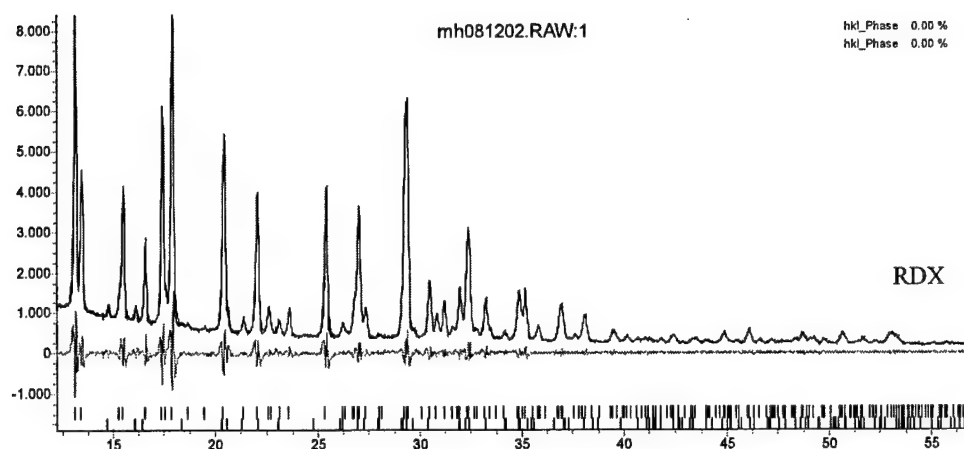


Fig. 6: Diffraction pattern and "hkl-phase fit" of the RDX-sample; positions of RDX and HMX peaks are marked by blue and black bars, respectively.

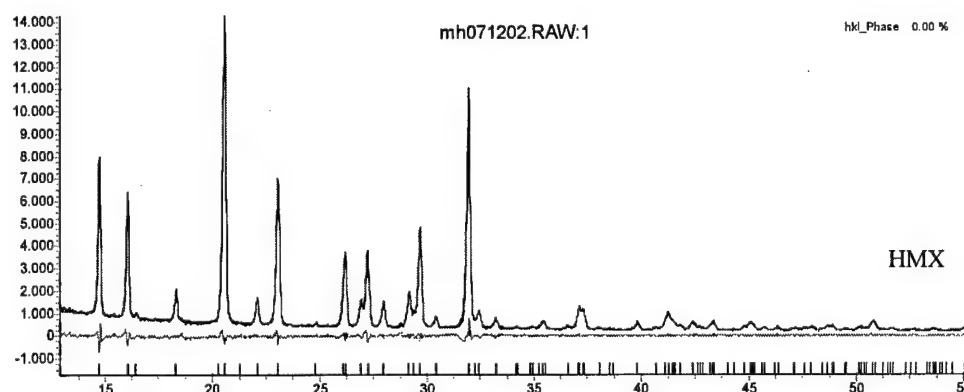


Fig. 7: Diffraction pattern and "hkl-phase fit" of the HMX-sample; all measured peaks belong to HMX.

Single peak fit

FWHM of Bragg-Brentano measurements / SPVII-profile

Fig. 8 till Fig. 10 show selected diffraction peaks of SRM 660a, RDX and HMX fitted by the SPVII analytical function. The calculated fitted well into the measured profiles; particularly, asymmetries of the RDX- and HMX-peaks are simulated accurately by the split function. Compared to the SRM 660a, the energetic materials yield strongly broadened and asymmetric peaks, particularly with common sample preparation.

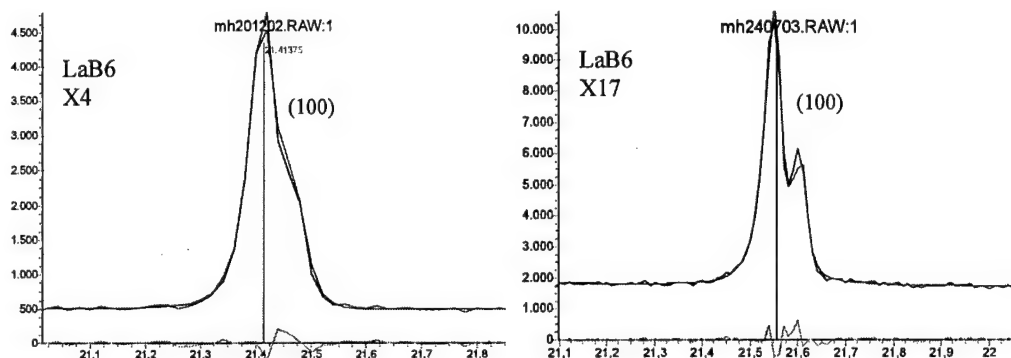


Fig. 8: Diffraction peaks of commonly prepared SRM 660a (left) and a powder layer (right) fitted by SPVII

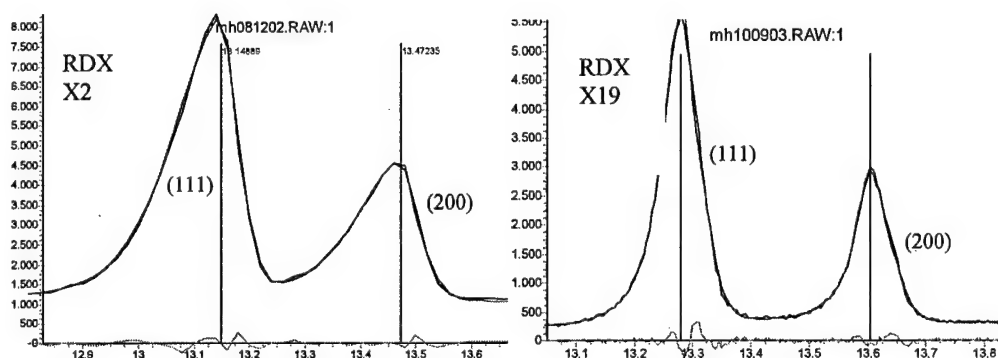


Fig. 9: Diffraction peaks of commonly prepared RDX (left) and a powder layer (right) fitted by SPVII

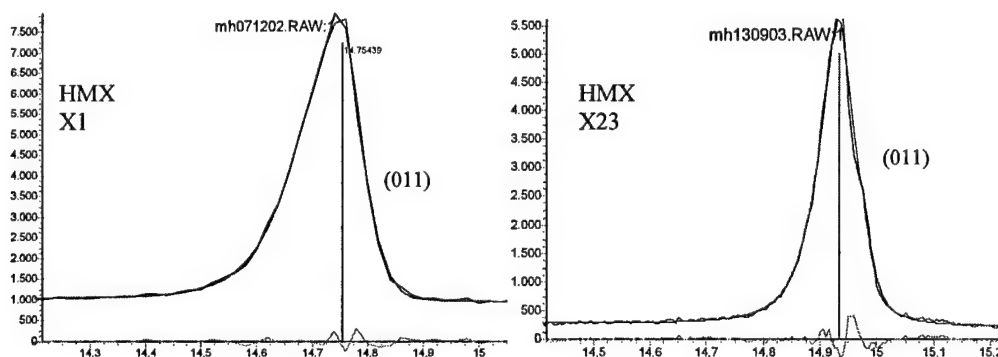


Fig. 10: Diffraction peaks of commonly prepared HMX (left) and a powder layers fitted by the split-Pearson VII analytical function (SPVII)

The fitted peak widths (FWHM) of samples measured with common sample preparation and with powder layers are plotted versus peak position in Fig. 11. It shows interesting details:

With the common preparation technique the peaks of the energetic materials are two to three times wider than those of the standard. The differences represent sample properties as absorption, crystallite size and micro strain.

Using the layer technique peaks of all samples slim down significantly. The effect increases from LaB6 to HMX to RDX, correlating with the calculated penetration depth in Tab. 2. The peak widths of

RDX are nearly as small as those measured with the standard LaB⁶, indicating that absorption broadening does not more hinder a proper profile analysis.

Comparing the results from powder layers, the peaks of HMX are wider than those of RDX, representing a larger size/strain broadening.

The results are reproducible as shown with the two RDX samples. Additionally, the plot reveals a raw agreement of the runs of the curves measured with both preparation techniques, which give a first idea about the unisotropy of sample properties.

No peaks of SRM 660a appear below 21.4° 2Theta (a reference material for the low 2Theta range should be used for a more accurate determination of geometric peak profiles and widths)

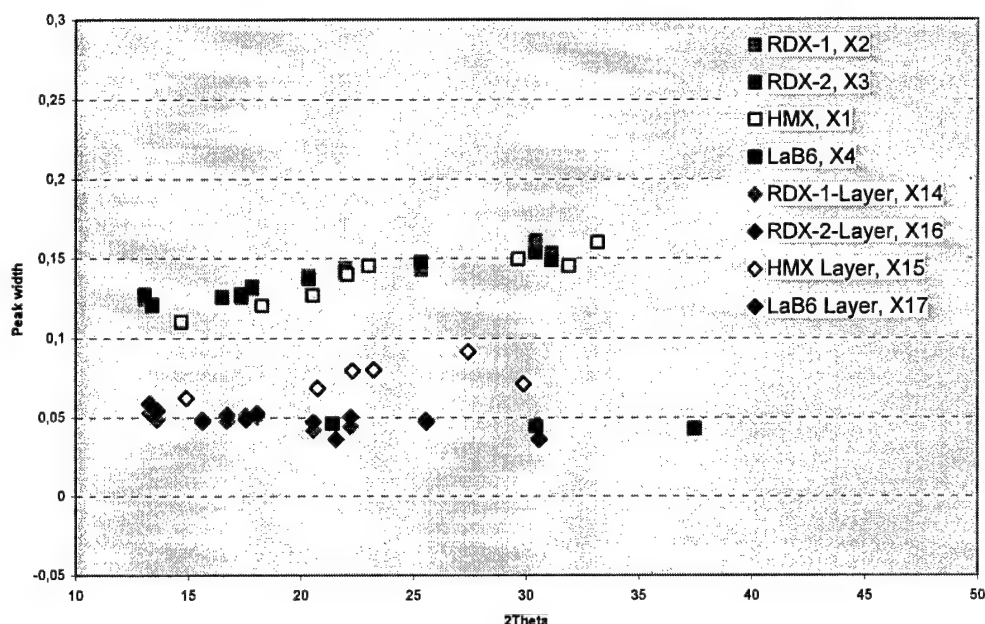


Fig. 11: FWHM of the single peaks of SRM 660a, RDX and HMX from common sample preparation and powder layers plotted versus 2Theta.

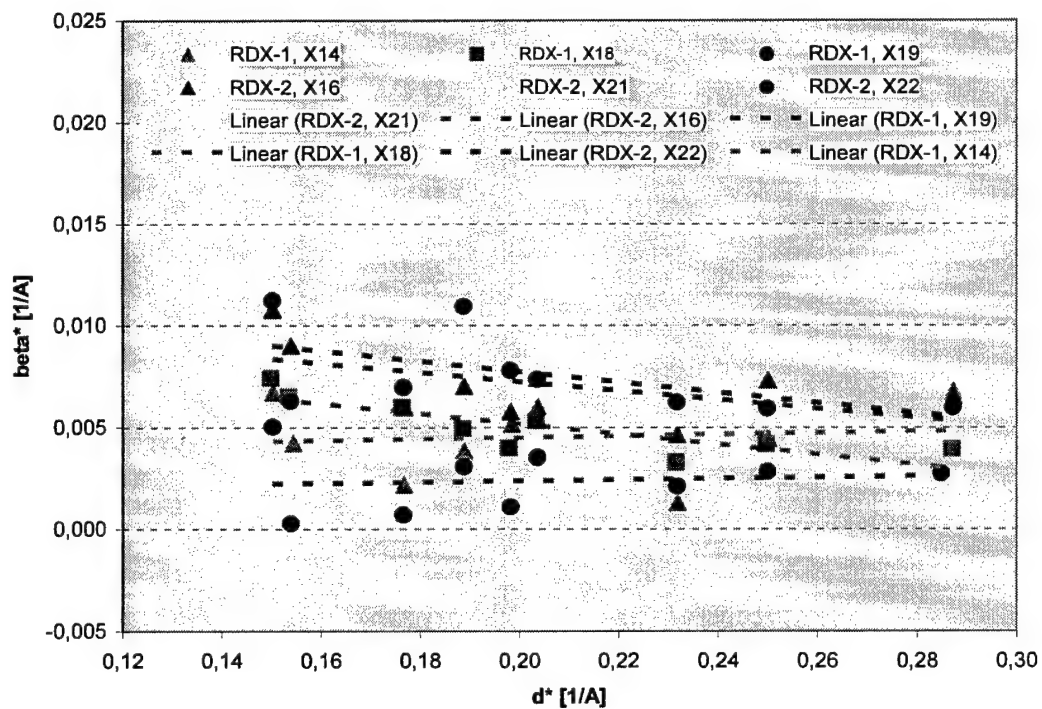


Fig. 12: Williamson-Hall-plot of RDX

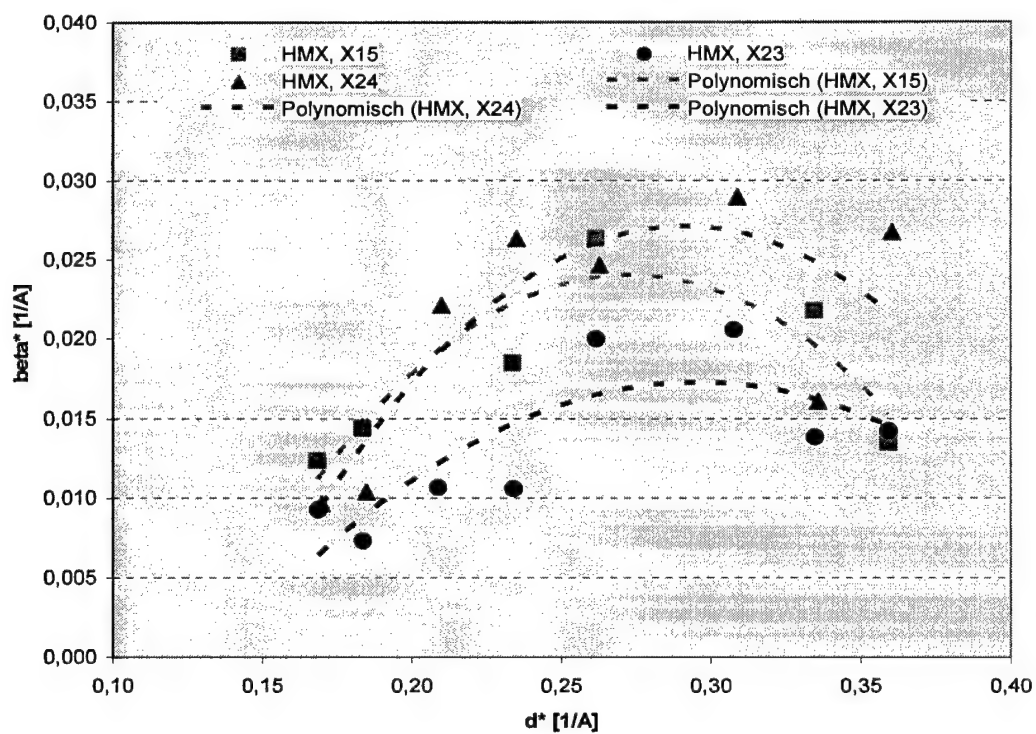


Fig. 13: Williamson-Hall-plot of HMX

Tab. 5: Regression line data from the Williamson-Hall-plots

$y=ax^2+bx+c$	a	b	c	R^2	Mean b
RDX-1, X14	0	0.0037	0.0038	0.008	-0.016(14)
RDX-1, X18	0	-0.0248	0.0101	0.66	
RDX-1, X19	0	-0.0255	0.0128	0.31	
RDX-2, X16	0	-0.0221	0.0117	0.3	-0.014(12)
RDX-2, X21	0	-0.0229	0.0159	0.38	
RDX-2, X22	0	0.003	0.0018	0.008	
HMX, X15	-1.1704	0.639	-0.0632	0.8	
HMX, X23	-0.6704	0.3965	-0.0414	0.64	
HMX, X24	-1.1883	0.6923	-0.0737	0.66	
< 0.33					0.127(19)
HMX, X15	0	0.1368	-0.0111	0.92	
HMX, X23	0	0.1001	-0.0097	0.82	
HMX, X24	0	0.1429	-0.0123	0.79	

Fig. 12 and Fig. 13 show Williamson-Hall-plots of RDX and HMX and their regression lines. The regression line parameters are summarized in Tab. 5.

The RDX data admit proper fits of linear regression lines with low or even negative slopes and more or less systematic deviations around the lines. In contrast the reciprocal peak widths of HMX at first increase strongly with increasing d^* , and fall down again beyond 0.33. Thus the HMX curves are better described by 2nd-order regression line, shown in Fig. 13. However, the initial slopes of the HMX-curves are well fitted by linear regressions in the d^* -range below 0.33. Yielded average slopes are -0.016, -0.014 and 0.127 for RDX-1, RDX-2 and HMX, respectively. The values indicate that the micro strain is much higher in HMX then in RDX, and it seems slightly higher in RDX-2 then in RDX-1.

The average deviations of the measured data from the linear and 2nd-order regression lines of RDX and for HMX, respectively, is shown in Fig. 14 and Fig. 15. The deviations of the RDX data are systematic and significant. Low values are found for the (hkl)-reflexes with $h=2$ and $l=0$, (200), (210) and (220), high values for (111), (002) and (102).

The deviations of the HMX data are less significant. It seems that (011) and (031) are strained and (020), (11-2) and (10-2) are unstrained peaks. However the low value of (022) contradict the high one of (011), and the standard deviations indicate that the values of (10-2), (022) and (13-2) are poorly determined. Further refined investigations of HMX seem to be necessary.

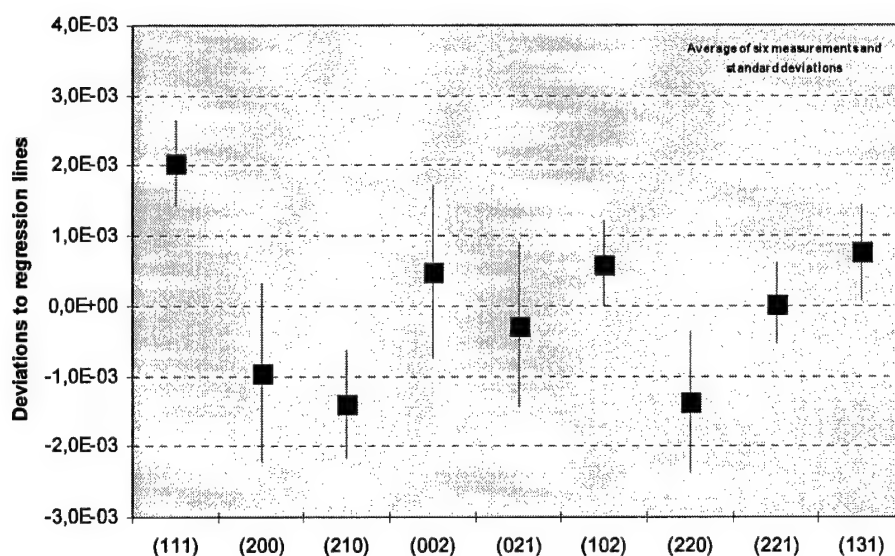


Fig. 14: Deviations of reciprocal peak widths from regression lines of RDX. Average values from six measurements.

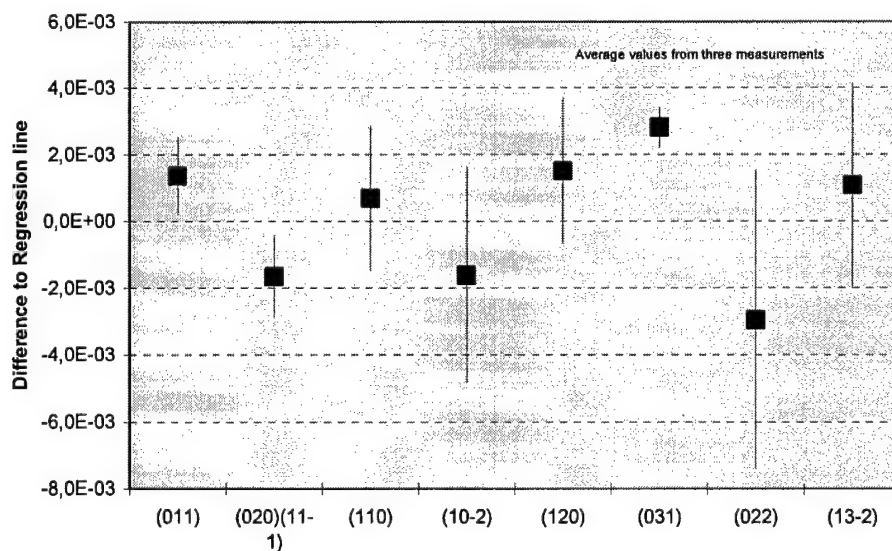


Fig. 15: Deviations of reciprocal peak widths from regression lines of HMX. Average values from three measurements.

Fundamental parameter approach FPA

The FPA fits as accurately as the SPVII-function. Tab. 6 summarizes the crystal size, micro strain and linear absorption coefficients evaluated from the three strongest peaks (standard deviations in brackets). The figures show a dependency of calculated crystal sizes and micro strain on the peak positions (2θ), particularly for RDX. Average particle sizes amount to 380, 417 and 667 μm at 13.3, 13.6 and 18.1 $^\circ 2\theta$ for RDX and to 273 μm for HMX, and the average micro strains amount

to 0.007, 0.006 and 0.0145 for RDX and 0.21 for HMX. The RDX-results of the measurements coincidence highly except for X21.

The results give rise for the hypothesis that the broader peaks measured with HMX stem from both, a lower crystal size and a higher micro strain. However, the accuracy of the method should be reinvestigated with further refined measurements.

Tab. 6: Crystal size, micro strain and linear absorption coefficients obtained with the fundamental parameter approach.

	2Theta	Aera	CrySize	MS	Abs.	Meas.
RDX	13,26	270	441	0,0001	40	X18
	13,29	508	405	0,0005	25	X19
	13,3	604	271	0,0271	126	X21
	13,28	151	404	0,0001	500	X22
	average		380(75)	0,007(13)		
	exc. X21		417(21)	0,0002(2)		
	13,59	122	466	0,0001	40	X18
	13,62	236	489	0,0005	25	X19
	13,63	290	275	0,0215	126	X21
	13,61	71	440	0,0001	500	X22
	average		417(97)	0,006(11)		
	exc. X21		465(24)	0,0002(2)		
	18,03	238	849	0,0192	25	X18
	18,05	461	602	0,0023	32	X19
	18,07	570	360	0,0202	61	X21
	18,06	142	858	0,0165	43	X22
	average		667(237)	0,0145(83)		
	exc. X21		770(145)	0,0127(90)		
HMX	14,95	513	481	0,0001	481	X23
	15,03	347	279	0,016	279	X24
	20,77	610	285	0,0045	285	X23
	20,88	374	156	0,0017	156	X24
	23,26	178	248	0,055	248	X23
	23,37	169	190	0,046	190	X24
	average		273(114)	0,021(24)		

Discussions

The medium particle size of the fine energetic materials, determined by SEM-investigations, lie between 3 to 5 μm , where no problems with sample preparation or poor statistical distribution should occur. For that reason no special X-ray optical devices are required for the measurements of the internal crystalline properties, in contrast to earlier investigations with coarse samples.

Consequently X-ray diffraction experiments yielded continuously shaped diffraction profiles, well suited for peak profile analysis. However, the diffraction peaks of the energetic materials were strongly broadened compared to those of the reference materials. Mayor parts of the peak broadening combined with strong asymmetries represent the low absorption of the nitramines with calculated penetration depths between 600 and 650 μm .

Preparing thin powder layers reduces the penetration depth and therefore the absorption broadening. Thus much narrower peaks until 0.04 $^{\circ}2\text{Theta}$ FWHM have been obtained, even for the LaB6 with its relatively strong absorption. A further advantage has to be considered in this context. With the

relatively low symmetry of the energetic materials and the resulting high quantity of appearing peaks in the patterns, most peaks or peak tails overlap with neighboring ones. The reduction of absorption broadening yields an improved peak separation, and therefore a more reliable peak profile analysis.

However, for the high resolution poor peak intensities or long measuring time, and a relatively high underground have to be accepted. Sample holders, with lower underground in the patterns and the measurement of reduced peak ranges around the interesting peaks may moderate these lacks. Besides, the makeshift preparation of powder layers with determined reproducible thickness and the positioning of the samples have to be refined, to increase the accuracy of the method.

Common and powder layer technique revealed significant differences between HMX and RDX. The evaluation with Williamson Hall-plot and with fundamental parameter approach, point to higher micro strain in HMX compared to RDX, and the fundamental parameter approach point to a smaller crystallite size in HMX compared to RDX. Considering the defect models described in literature, dislocations in RDX and twins in HMX, it seems that twins have a stronger effect on peak broadening than dislocations. This could be discussed in terms of the different features of these defect types. Strained areas of twins are related to interfaces, whereas areas of dislocations are related to screw or edge like lines. The two-dimensional character of twins may strain a higher portion of crystals than the one dimensional of dislocations. Besides, twin interfaces separate domains and therefore may reduce the size of coherently diffracting crystallites.

A very interesting detail is the significant unisotropic peak broadening, particularly measured with the powder layer technique. The RDX peaks (111), (002) and (102) are broadened compared to (200), (210) and (220). Considering edge dislocations, broadening would occur at peaks representing net planes parallel to the edge of a dislocation, but not perpendicular. Starting with low indices [100] or [110] could represent the edge of a dislocation and [001] its burger vector. In HMX the peaks (011) and (031) are broadened compared to (020), (11-2) and (10-2). Considering twinning in HMX this could point to (011) twin planes or to unisotropic particle formation during crystallization. However, the unisotropic broadening in HMX seems to be less significant than those observed in RDX, and the results give an incomplete idea on line broadening, as only a few peaks could be separated and evaluated. No information was obtained about peaks characterizing the models given in Literature, e.g. the [-100](040) edge dislocation in RDX and (101)[10-1] twinning in HMX.

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Any opinion, findings and conclusions or recommendations expressed in this material are those of the authors and do not necessarily reflect the view of the European Office of Aerospace Research and Development, Air Force Office of Scientific Research, Air Force Research Laboratory.

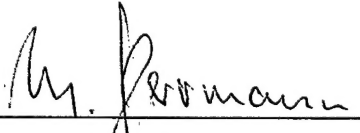
Declarations

The Contractor, FRAUNHOFER INSTITUT FÜR CHEMISCHE TECHNOLOGIE., hereby declares that, to the best of his knowledge and belief, the technical data delivered herewith under Contract No. FA8655-02-M4066 is complete, accurate, and complies with all requirements of the contract.

I declare that there were no subject inventions to be declared as defined in FAR 52.227-13, during the performance of this contract.

Date:

16.12.03


(Dr. Michael Herrmann)

Appendix

Tab. 7: Diffraction measurements and conditions

#	Sample	System/Sample	2Theta/Step	Time/Ch.	Rotation	File
X1	HMX	BB	5-90/0.02	8 s	30 rpm	Mh071202
X2	RDX-1	BB	5-90/0.02	8 s	30 rpm	Mh081203
X3	RDX-2	BB	5-90/0.02	8 s	30 rpm	Mh091203
X4	LaB6	BB	5-90/0.02	8 s	30 rpm	Mh201203
X5	RDX-1	DS*-Capillary	10-40/0.01			Mh040603
X6	HMX	DS-Capillary	10-40/0.01			Mh050603
X7	RDX-2	DS-Capillary	10-40/0.01			Mh060603
X8	LaB6	DS-Capillary	10-40/0.01			Mh070603
X9	HMX	BB	10-35/0.01	10 s	30 rpm	Mh160703
X10	RDX-2	BB	10-35/0.01	10 s	30 rpm	Mh170703
X11	RDX1	BB	10-35/0.01	10 s	30 rpm	Mh180703
X12	RDX-1	BB-Wafer	10-35/0.01	10 s		Mh190703
X13	RDX-1	BB-Wafer	10-35/0.01	10 s	30 rpm	Mh200703
X14	RDX-1	BB-Alu	10-35/0.01	10 s	30 rpm	Mh210703
X15	HMX	BB-Alu	10-35/0.01	10 s	30 rpm	Mh220703
X16	RDX-2	BB-Alu	10-35/0.01	10 s	30 rpm	Mh230703
X17	LaB6	BB-Alu	10-35/0.01	10 s	30 rpm	Mh240703
X18	RDX-1	BB-Alu	12,5-26/0.005	10 s	30 rpm	Mh090903
X19	RDX-1	BB-Alu	12,5-26/0.005	10 s	30 rpm	Mh100903
X21	RDX-2	BB-Alu	12,5-26/0.005	10 s	30 rpm	mh110903
X22	RDX-2	BB-Alu	12,5-26/0.005	10 s	30 rpm	mh120903
X23	HMX	BB-Alu	14-33/0.005	10 s	30 rpm	mh130903
X24	HMX	BB-Alu	14-33/0.005	10 s	30 rpm	mh140903
X25	LaB6	BB-Alu	14-33/0.005	10 s	30 rpm	mh150903

*BB=Bragg-Brentano, DS=Debye-Scherrer

Tab. 8: Micro strain and crystal size of RDX and HMX obtained by fundamental parameter approach measured with common sample preparation

RDX				HMX			
Position [2Theta]	hkl	Micro strain	Absorption [1/cm]	Position [2Theta]	hkl	Cry Size L(nm)	Absorption [1/cm]
13.17	111	0.0289	4.1	14.77	011	960	5.3
13.49	200	0.0027	4.1	18.35	110	360	6.9
16.61	002	0.0104	5.1	20.60	10-2	352	7.1
17.44	021	0.0304	5.1	22.13	11-2	175	7.8
17.93	102	0.0355	5.1	23.09	120	239	7.8
20.44	220	0.0808	6	27.27	031	240	6.8
22.08	221	0.0571	6	29.73	022	460	7.0
25.44	131	0.0301	6.5	31.98	13-2	457	9.0
30.51	322	0.0917	7.8				

Tab. 9: Indexing (hkl) by Rietveld Analysis and selection of separated peaks (yellow) of RDX and HMX.

RDX						HMX					
h	k	l	d	2Theta	int.	h	k	l	d	2Theta	int.
1	1	1	6,76777	13,071	1005	0	1	1	6,02244	14,697	916
2	0	0	6,60217	13,400	466	1	1	-1	5,5278	16,020	321
0	2	0	5,80761	15,244	64	0	2	0	5,52402	16,031	445
2	1	0	5,73974	15,425	447	1	0	0	5,39711	16,411	9
0	0	2	5,36515	16,509	290	1	1	0	4,8494	18,280	183
0	2	1	5,10751	17,349	818	0	2	1	4,37901	20,263	48
2	1	1	5,06116	17,509	60	1	0	-2	4,32364	20,525	2207
1	0	2	4,97052	17,831	1155	1	2	-1	4,1774	21,252	0
1	2	1	4,76357	18,612	0	1	1	-2	4,0263	22,059	175
1	1	2	4,56968	19,409	0	1	2	0	3,86042	23,020	1099
2	2	0	4,3606	20,349	821	0	0	2	3,59179	24,768	3
2	0	2	4,16369	21,323	48	0	1	2	3,41581	26,066	157
2	2	1	4,03977	21,985	590	1	2	-2	3,40474	26,152	563
0	2	2	3,94088	22,544	119	1	1	1	3,31357	26,885	238
2	1	2	3,91948	22,668	11	0	3	1	3,27714	27,189	653
3	1	1	3,84285	23,127	49	2	0	-2	3,1922	27,927	137
1	2	2	3,77628	23,540	109	1	3	-1	3,19002	27,947	100
1	3	1	3,51082	25,348	705	2	1	-2	3,06676	29,094	303
3	0	2	3,40287	26,167	76	2	1	-1	3,06366	29,124	34
2	2	2	3,38389	26,316	30	1	3	0	3,04199	29,336	106
2	3	0	3,33981	26,670	0	0	2	2	3,01122	29,643	910
3	2	1	3,33421	26,715	155	1	2	1	2,94048	30,373	119
1	1	3	3,30927	26,920	527	1	3	-2	2,80355	31,895	2243
4	0	0	3,30108	26,988	209	1	1	-3	2,77989	32,174	0
3	1	2	3,26561	27,287	153	2	2	-2	2,7639	32,365	70
2	3	1	3,18891	27,957	34	0	4	0	2,76201	32,388	66
4	1	0	3,17534	28,079	25	2	2	-1	2,76163	32,393	38
1	3	2	3,05444	29,214	866	2	0	0	2,69856	33,171	145
0	2	3	3,04551	29,302	335	2	1	-3	2,62732	34,098	26
4	1	1	3,04482	29,309	565	2	1	0	2,62149	34,176	27
2	1	3	3,0356	29,400	0	0	4	1	2,57802	34,770	26
1	2	3	2,9676	30,089	43	0	3	2	2,57131	34,864	33
3	2	2	2,936	30,421	386	1	2	-3	2,54839	35,188	36
0	4	0	2,9038	30,766	168	1	4	-1	2,53496	35,380	95
4	2	0	2,86987	31,139	266	1	3	1	2,52684	35,498	31
2	3	2	2,83533	31,528	92	1	4	0	2,45875	36,515	72
4	0	2	2,81152	31,802	11	2	2	-3	2,42932	36,973	48
3	3	1	2,80592	31,868	157	2	2	0	2,4247	37,046	185
0	4	1	2,80298	31,902	188	1	0	2	2,42397	37,058	24
4	2	1	2,77243	32,263	516	2	3	-2	2,41214	37,246	104
2	2	3	2,76547	32,346	339	2	3	-1	2,41063	37,271	119
1	4	1	2,74188	32,632	0	1	1	2	2,36766	37,973	52
4	1	2	2,73261	32,746	83	0	1	3	2,34019	38,436	34
3	1	3	2,69977	33,156	302	1	4	-2	2,32761	38,652	27
0	0	4	2,68258	33,375	70	1	3	-3	2,26487	39,767	138
2	4	0	2,65807	33,692	59	1	2	2	2,21967	40,612	62
1	0	4	2,62887	34,077	122	0	2	3	2,197	41,050	83
2	4	1	2,58008	34,742	192	0	4	2	2,1895	41,197	205